



Cite this: *J. Mater. Chem. C*, 2021,
9, 10889

Enhanced electronic communication through a conjugated bridge in a porphyrin–fullerene donor–acceptor couple†‡

Rubén Caballero, ^a Joaquín Calbo, ^b Juan Aragó, ^b Pilar de la Cruz, ^a Enrique Ortí, ^{*b} Nikolai V. Tkachenko ^{*c} and Fernando Langa ^{*a}

A ZnP–2EDOTV–C₆₀ triad, with enhanced electronic communication between terminus donor and acceptor moieties, was synthesized and studied both experimentally and theoretically. Electrochemical measurements and density functional theory calculations support that the first oxidation takes place on the 3,4-ethylenedioxythiophenevinylene (2EDOTV) bridge followed by the oxidation of the ZnP moiety at slightly higher energies. The electronic communication between the terminal electron-donor ZnP and the electron-acceptor C₆₀ units is enhanced by the conjugated EDOTV-based spacer leading to photo-induced electron transfer over the distance >2 nm in the picosecond time domain. The involvement of the spacer in the electron transfer is confirmed by both photophysical measurements and theoretical calculations indicating the formation of intermediate ZnP–2EDOTV⁰⁺–C₆₀^{δ-} charge-transfer states. Overall, the enhanced donor–acceptor electronic communication accelerates the photoinduced charge separation but also affects the charge recombination process.

Received 12th March 2021,
Accepted 25th June 2021

DOI: 10.1039/d1tc01160j

rsc.li/materials-c

Introduction

The challenge of using renewable energies instead of fossil fuels to fight the global warming is one of the main scientific tasks for the next few decades and the utilization of sunlight is the most promising alternative.^{1,2} A general approach is to design a system in which the energy of an excited state is converted to the energy of an electron–hole pair, or a cation–anion radical pair in the case of molecular system, which is used to generate electric current or to initiate a chemical reaction, respectively. In either case, it is important to achieve an as-long-as-possible charge separation (CS) to avoid fast charge recombination and allow efficient utilization of the charges or ions. In natural photosynthesis, this is achieved through multiple electron transfer reactions. The reaction centre consists of a series of redox active molecular entities

placed close to each other such that the excitation of a primary electron donor (D) triggers a cascade of electron transfer (ET) reactions delivering an electron to a final electron acceptor (A) at a few nanometer distance from the primary donor.^{3–5} Natural photosynthesis utilizes chlorophylls and quinones as essential components,⁶ but numerous artificial systems were developed utilising porphyrins, the closest analogues of chlorophylls, and fullerenes, which are equivalent to quinones in their reduction potentials but have the advantage of a smaller reorganization energy associated with the ET.^{7–10} Although this strategy allows to design systems with almost unity quantum yield of the CS, it suffers from energy loss at each step of the ET and cumulative loss of energy from the excited state to the final CS state can be high, more than 50%. The ET distance for a single step of so-called through-space ET is limited by the fact that the electron must “jump” from a donor to an acceptor, and typically an increase of the distance by 1 Å reduces the ET time constant by factor of two.¹¹

An alternative approach is to design donor–bridge–acceptor (D–B–A) systems, in which the bridge plays the role of a molecular wire conducting an electron from the lowest-unoccupied molecular orbital (LUMO) of the donor to that of the acceptor.^{12,13} This requires that the LUMO of the bridge is located close to that of the donor, which decreases the ET rate sensitivity to the D–A distance by an order in magnitude or even more.^{13,14} The connection between the donor and acceptor, the bridge, therefore becomes an important player in the

^a Instituto de Nanociencia, Nanotecnología y Materiales Moleculares (INAMOL), Universidad de Castilla-La Mancha, Campus de la Fábrica de Armas, 45071 Toledo, Spain. E-mail: fernando.langa@uclm.es

^b Instituto de Ciencia Molecular, Universidad de Valencia, 46950 Paterna, Spain. E-mail: enrique.orti@uv.es

^c Faculty of Engineering and Natural Science, Tampere University, P. O. Box 541, Tampere 33101, Finland. E-mail: nikolai.tkachenko@tuni.fi

† Dedicated to Prof. Jaume Veciana and Prof. Concepció Rovira on occasion of their 70 birthdays.

‡ Electronic supplementary information (ESI) available. See DOI: 10.1039/d1tc01160j

photoinduced ET of these systems. From this viewpoint, the design of efficient D-B-A systems required a careful adjustment of energies of all the three components in conjunction with each other. This in turn rises the requirements to the chemical engineering since the system cannot be considered as consisting of two relatively independent parts, the donor and the acceptor, but all three components must work coherently. Though this is a complication, the expected outcome is the longer distance of the CS with smaller energy losses.

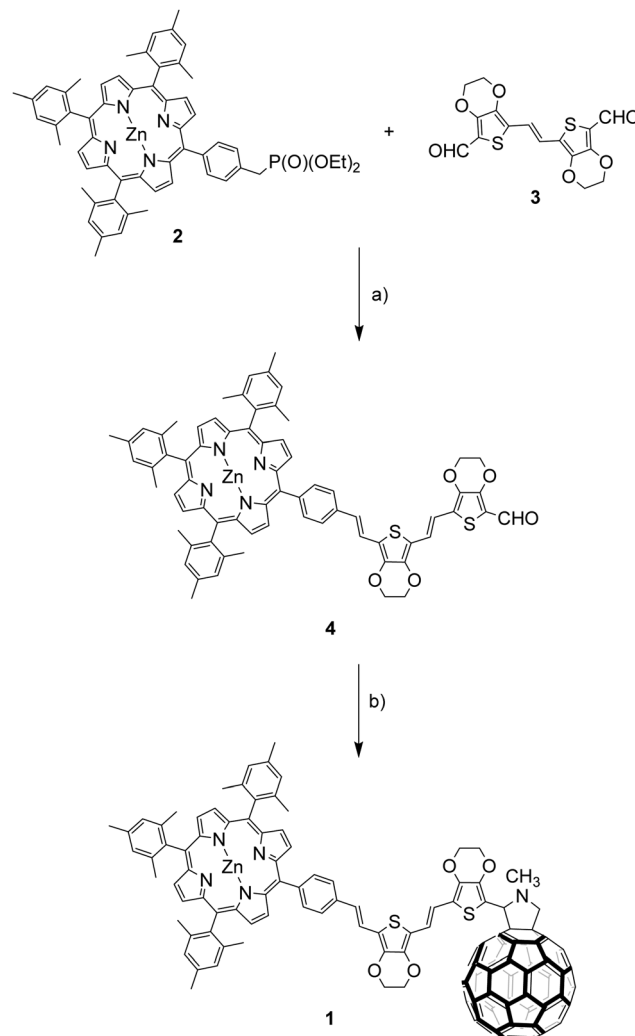
To this end, porphyrins and fullerenes have been widely used as D and A, respectively, in D-B-A systems,^{15,16} whereas conjugated oligomers have been employed as bridges because they play an important role as semiconductors in molecular electronics.¹⁷ Porphyrin-B-C₆₀ architectures using *p*-phenylenevinylene,¹⁸ *p*-phenylene ethynylene,¹⁹ fluorene,²⁰ thiophene²¹ or thiylenevinylene^{10,22} oligomers as bridges have been shown to exhibit efficient long-distance photoinduced electron transfer from the porphyrin to the fullerene.

In this work, the synthesis of the donor-spacer-acceptor ZnP-2EDOTV-C₆₀ (**1**) triad, composed by a Zn-porphyrin as donor, a π -conjugated rigid oligomer formed by two units of 3,4-ethylenedioxothiophenevinylene (2EDOTV) as bridge and [60]fulleropyrrolidine as acceptor, is reported. Oligothienylenevinylanes (nTVs) have been reported to be one of the most planar and versatile conjugated oligomers described so far with a low attenuation factor.^{10,22,23} In addition, the inclusion of ethylenedioxy groups allows to obtain nEDOTVs, which, compared with nTVs, stabilize more efficiently positive charges.²⁴ The electrochemical and photophysical properties of the new triad are studied in depth to get insight into the influence that the electronic coupling between the terminal **ZnP** donor and C₆₀ acceptor units, through the 2EDOTV conjugated bridge, has over the photoinduced electron transfer. Theoretical calculations at the DFT level, including solvent effects, and detailed femtosecond transient absorption spectroscopic studies have been performed to investigate the electron transfer process.

Results and discussion

Synthesis

The synthesis of triad ZnP-2EDOTV-C₆₀ (**1**) was achieved following a convergent methodology (Scheme 1). First, porphyrin **2**²⁵ and bisaldehyde **3**²⁶ were reacted by Wittig–Horner olefination, under careful stoichiometric control, affording **4** in 42% yield. The ¹H-NMR spectrum of **4** shows the H atom of the aldehyde at 9.83 ppm as a singlet (Fig. S1, ESI[‡]). Next, **1** was obtained by 1,3-dipolar cycloaddition of C₆₀ and the azomethyne ylide generated *in situ* from **2** and *N*-methylglycine. The target compound **1** was isolated by column chromatography (silica gel, hexane : toluene 1 : 1), followed by gel permeation chromatography (Biobeads S-X1, toluene) to obtain a purple solid (44% yield). The MALDI-TOF MS spectrum shows the correct molecular ion at *m/z* = 1910.69 amu with a fragmentation peak at *m/z* = 720 amu due to the fullerene cage (Fig. S8, ESI[‡]). Triad **1** was characterized by ¹H-NMR spectroscopy showing the signals corresponding to



Scheme 1 Synthesis of triad ZnP-2EDOTV-C₆₀ (**1**). *Reagents and conditions:* (a) NaH in THF at 0 °C. (b) C₆₀ and *N*-methylglycine in chlorobenzene at reflux.

the porphyrin macrocycle between 8.7 and 9.0 ppm, and the vinyl signals of the oligomer as an AA'BB' system between 6.9 and 7.5 ppm (Fig. S5, ESI[‡]). All double bonds are obtained as the *E* isomer as proved by the coupling constants (*J* = 16 and 16.2 Hz). The fingerprint of the fulleropyrrolidine ring appears as a singlet at 5.43 ppm and two doublets at 4.98 and 4.23 ppm. Synthetic details and NMR, FT-IR and MALDI-TOF MS data for molecules **1** and **4** are given in the Experimental section included in the ESI[‡].

In addition, compounds **5** (Ph-2EDOTV), **6** (Ph-2EDOTV-C₆₀), Zinc(II) 5,10,15,20-tetrakis(2,4,6-trimethylphenyl)porphyrin (**ZnP**) and *N*-methyl-[60]fulleropyrrolidine (**NMFP**) (Chart 1) were prepared as reference systems for comparison purposes (see the ESI[‡]).

Geometry and molecular orbitals

Theoretical calculations under the density functional theory (DFT) framework were performed for geometry and electronic structure analysis (see the ESI[‡] for full computational details). The minimum-energy structure of triad **1** (Fig. 1a) and its

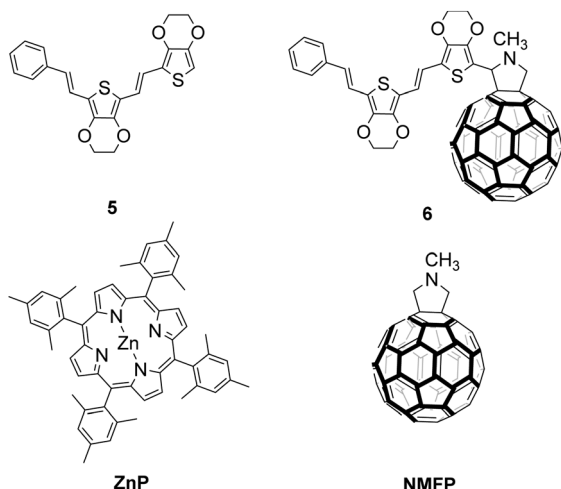


Chart 1 Reference compounds Ph-2EDOTV (**5**), Ph-2EDOTV-C₆₀ (**6**), ZnP and NMFP

constituting derivatives **4**, **5**, **6**, **ZnP** and **NMFP** (Fig. S26, ESI[‡]) were obtained after full geometry relaxation at the B3LYP/cc-pVDZ+LANL2DZ(Zn) level of theory including solvent effects (dichloromethane in a polarizable continuum model PCM approach). Triad ZnP-2EDOTV-C₆₀ (**1**) shows a non-coplanar

arrangement of the three constituting moieties (porphyrin, bridge and fullerene) with an almost orthogonal disposition (70°) of the π -conjugated 2EDOTV bridge with respect to the **ZnP** unit due to the internal rotation of the *meso*-phenyl ring (Fig. 1a). Among the different possible conformations of the 2EDOTV bridge, the one containing *E* double bonds with an *anti* orientation of the EDOT groups and sulphur atoms pointing inwards in the π -conjugated chain is found as the most stable arrangement (Fig. S26, ESI ‡).

Electronic structure calculations were performed on the minimum-energy geometries to unveil the nature of the frontier molecular orbitals in triad **1** (Fig. 1b) and relative compounds (Fig. S27 and S28, ESI[‡]). The highest-occupied molecular orbital (HOMO) of **1** is calculated at -4.83 eV and spreads over the π -conjugated bridge including the *meso*-phenyl ring (Fig. 1b). The HOMO shows a topology and an energy similar to those computed for the HOMOs of related Ph-2EDOTV (5), Ph-2EDOTV-C₆₀ (6) and ZnP-2EDOTV (4) systems (Fig. S27 and S28, ESI[‡]). In contrast, the HOMO-1 and HOMO-2 of **1** are calculated at -5.21 and -5.43 eV, respectively, and are localized over the porphyrin moiety, in good analogy to that predicted for **4** and to the respective HOMO and HOMO-1 of **ZnP** (Fig. S27 and S28, ESI[‡]). These results suggest that the 2EDOTV bridge is a slightly stronger electron donor system compared to **ZnP**. The lowest-unoccupied molecular orbitals (LUMO, LUMO+1 and LUMO+2) of triad **1** are of C₆₀ nature, in accordance with those computed for **6** and also for **NMFP** (Fig. S27, ESI[‡]), indicating the acceptor nature of the fullerene moiety. For **4**, due to the absence of the C₆₀ moiety, the LUMO is calculated high in energy at -2.59 eV and belongs to the 2EDOTV bridge, whereas the LUMO+1 and LUMO+2 are placed over the porphyrin unit (Fig. S27, ESI[‡]).

Electrochemical properties

The redox behaviour of triad **1** and of the reference compounds **ZnP**, **NMFP**, **5** and **6** was investigated by Oyster-Young square-wave voltammetry (OSWV), and the reversibility of each redox process was checked by cyclic voltammetry (CV). Table 1 collects the values measured for the first redox potentials.

A complex cyclic voltammogram showing several reduction and oxidation processes was obtained for **1** (Fig. S33, ESI[‡]). The contribution of each electroactive moiety of the triad can be addressed by comparing the redox potentials measured for **1**

Table 1 Redox potentials (V vs. Fc/Fc⁺) of the processes observed by OSWV for compounds **1**, **5**, **6**, **ZnP** and **NMFP**^a

	E_{red}^2	E_{red}^1	E_{ox}^1	E_{ox}^2
ZnP	-2.55	-2.02	0.25	0.65
NMFP	-1.59	-1.16		
5		-2.37 ^b	0.16 ^b	
6	-1.55	-1.15	0.19 ^b	
1	-1.55	-1.15	0.21	0.28

^a 10⁻³ M in *o*-dichlorobenzene:acetonitrile (4:1), Ag/AgNO₃ (0.01 M) was used as reference electrode and checked against the (Fc/Fc⁺) couple, glassy carbon electrode, Pt counter electrode, 20 °C, 0.1 M Bu₄NClO₄, scan rate = 100 mV s⁻¹. ^b Irreversible process.

Fig. 1 (a) Minimum-energy structure (characteristic dihedral angles are indicated) and (b) frontier molecular orbital contours (isovalue = ± 0.03) calculated for triad **1** at the B3LYP/cc-pVDZ+LANL2DZ(Zn) level including solvent effects. H and L denote HOMO and LUMO, respectively.

with those of the reference compounds (Table 1). The reduction potentials of **1**, registered at -1.15 and -1.55 V, are unequivocally assigned to the reduction of the C_{60} cage by comparison with those of **NMFP** and **6**. In the anodic window, several waves corresponding to both the porphyrin moiety and the bridge were recorded. The first oxidation is attributed to the overlap of two waves at 0.21 and 0.28 V as can be deduced by careful deconvolution of the Oyster-Young voltammogram (Fig. S34, ESI[‡]). The similarity of the first oxidation potential of **1** (0.21 V) with that of **5** (0.16 V) and **6** (0.19 V) suggests that this process is centred on the oligomer bridge. Indeed, the shift to higher potentials of the oxidation of oligothienylenevinylenes linked to electron acceptors has been reported, and it is attributed to the presence of different rotameric conformations of the oligomer unit when attached (or not) to the bulky C_{60} .²⁷ The second oxidation peak of **1**, at 0.28 V, is then attributed to the porphyrin macrocycle.

From these data, the energy of the radical ion pairs (E_{RIP})²⁸ for $ZnP-2EDOTV^{+}-C_{60}^{-}$ and $ZnP^{+}-2EDOTV-C_{60}^{-}$ can be estimated as 1.36 and 1.44 V, respectively. These results suggest that the energy of both radical ion pairs is similar and that, in the eventual formation of a radical ion pair, both transient species would have almost the same energy.

Based on the results of the redox potential measurements, the energy diagram of HOMO and LUMO level relative positions can be drawn as presented in Fig. 2, though evaluations of the energies of the charge-separated (CS) states in the diagram do not account for the Coulomb potentials and can be taken as rough estimations only.

Theoretical calculations were performed on the charged species of triad **1** and reference compounds at the UB3LYP/cc-pVQZ+LANL2DZ(Zn)+PCM(dichloromethane) level to confirm the nature of the electrochemical processes recorded experimentally. Electron injection in the first and second reductions of **1** occurs on the electron-acceptor fullerene moiety, as indicated by the Mulliken net charge accumulated in C_{60} of $-1.01e$ for the anion and of $-1.99e$ for the dianion, to

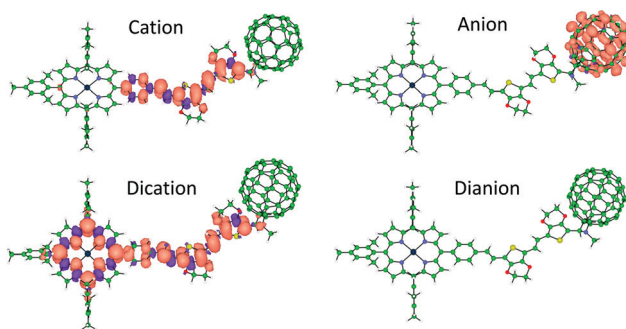


Fig. 3 Spin densities calculated for the lowest-energy spin-configuration of the singly and doubly oxidized and reduced species of triad **1**. Cation and anion: doublet state, dication: triplet state, dianion: closed-shell singlet state.

be compared with $-0.04e$ in the neutral species (Table S1, ESI[‡]). These processes are predicted with a first and second adiabatic electron affinity of 3.34 and 2.62 eV, respectively. As shown in Fig. 3, the electron-unpaired spin-density contours computed for the anion and dianion species of **1** further support the exclusive participation of C_{60} in the first and second reductions. First ionization energies (IE1) calculated for the cation species indicate almost identical values for triad **1** and referable compounds **5** and **6** (IE1 of 4.67 , 4.69 and 4.68 eV, respectively), in good correlation with HOMO energies (Fig. S28, ESI[‡]), suggesting a complete participation of the 2EDOTV bridge in the first oxidation process. On the other hand, the second oxidation of **1** is calculated with an IE2 = 5.27 eV, similar to the IE1 computed for referable **ZnP** (5.11 eV), indicating a main role of the porphyrin moiety in this oxidation process. Spin densities (Fig. 3), as well as accumulated Mulliken net charges (Table S1, ESI[‡]), further support that the first oxidation process takes place on the 2EDOTV bridge (cation in Fig. 3), whereas the second electron extraction occurs on the porphyrin unit (dication in Fig. 3), thus confirming the experimental assignments.

Absorption and emission properties

To get insight into the optical properties of the D-B-A triad **1**, the absorption spectrum of **1** and **ZnP** were recorded in CH_2Cl_2 and are shown in Fig. 4, whereas those of compounds **5** and **6** are displayed in Fig. S35 (ESI[‡]). Model compound Ph-2EDOTV (**5**) shows a structured absorption band with maximum at 424 nm, assigned to a $\pi-\pi^*$ transition, which red-shifts up to 435 nm in **6** owing to the interaction with the C_{60} cage. On the other hand, **ZnP** shows the characteristic Soret- and Q-band absorptions at 419 nm and 551 and 591 nm, respectively. The spectrum of compound **1** is the sum of the spectra of the three molecular components. Besides the characteristic broad absorption of the fullerene cage in the $300-400$ nm region, the Soret band of the porphyrin moiety decreases in intensity and is slightly red-shifted (2 nm) and broadened compared to **ZnP**, probably due to the lower symmetry of the porphyrin macrocycle (Fig. 4). At lower energies, a broad band associated to the $\pi-\pi^*$ transition of the conjugated bridge is observed in

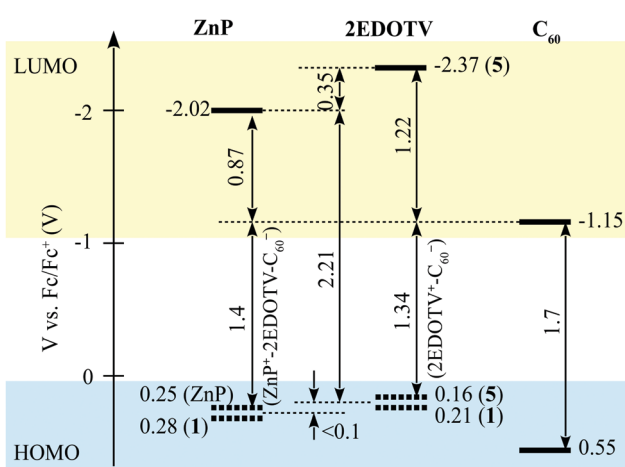


Fig. 2 Relative energy positions of HOMO and LUMO levels of the chromophores constituting the $Zn-2EDOTV-C_{60}$ triad.

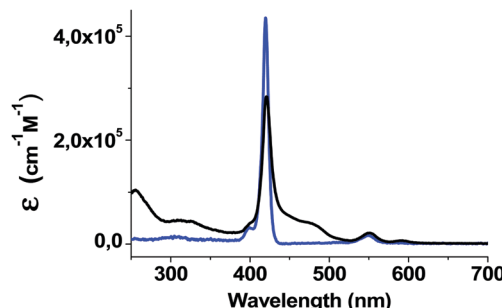


Fig. 4 UV-visible absorption spectra of ZnP-2EDOTV-C₆₀ (**1**, black) and ZnP (blue) in dichloromethane solution. Concentration: 10⁻⁵ M.

the 450–500 nm region, and the components of the Q-band of **ZnP** appear at similar energies.

The absorption spectra of **1**, **ZnP**, **5** and **6** were calculated at the TD-B3LYP/(cc-pVQZ+LANL2DZ(Zn))+PCM level using dichloromethane as solvent. Theoretical calculations predict the Q band of **ZnP** at 546 nm resulting from the S₀ → S₁, S₂ degenerate excitations, which exhibit a small oscillator strength ($f = 0.016$) and are due to π - π^* transitions centred over the porphyrin core (Fig. S36, ESI[‡]). The Soret band is predicted at 400 nm and is associated to a very intense doubly degenerate transition (S₀ → S₃, S₄, $f = 1.601$ each) of π - π^* nature. For **5** (Fig. S37, ESI[‡]), the S₀ → S₁ π - π^* transition is calculated at 499 nm with high intensity ($f = 1.491$), slightly underestimating the energy of the experimental band around 450 nm (Fig. S35, ESI[‡]). This band red shifts to 516 nm in **6**, in good agreement with the experiment. When moving to triad **1**, several low-lying charge-transfer excitations are predicted in the low-energy range (Fig. S38 and Table S2, ESI[‡]). The first S₀ → S₁, S₂, S₃ transitions are calculated at 981, 902 and 765 nm, respectively, with small or negligible oscillator strengths ($f = 0.002$, 0.032 and 0.000, respectively). These transitions are described by monoelectronic excitations from the HOMO (2EDOTV) to the LUMO, LUMO+1 and LUMO+2 (C₆₀), thus implying a charge transfer from the bridge to the fullerene ball (Fig. 1). Transitions S₀ → S₄, S₅, ..., S₁₀ computed below 700 nm mainly correspond to excitations from the porphyrin moiety to the C₆₀ cage and present a negligible intensity ($f < 0.001$) due to the zero overlap between the molecular orbitals of the **ZnP** and C₆₀ moieties (porphyrin-C₆₀ centroid-centroid distance of 24.5 Å). Then, an intense CT transition from the 2EDOTV bridge to the porphyrin is predicted at 591 nm (S₀ → S₁₁, $f = 0.767$), and the characteristic π - π^* excitation of the 2EDOTV bridge is calculated at 523 nm (S₀ → S₂₇, $f = 1.520$). These two intense transitions are calculated too low in energy and presumably give rise to the intense broad shoulder experimentally recorded in the 450–500 nm range (Fig. 4). The characteristic Q and Soret bands of the porphyrin moiety in triad **1** are predicted at similar energies as those calculated for **ZnP** in good agreement with the experimental data. The Soret band originates from transitions S₀ → S₅₃ (404 nm, $f = 0.830$) and S₀ → S₅₇ (399 nm, $f = 1.121$), which display lower intensities than for **ZnP** ($f = 1.601$) in good accord with the intensity decrease observed

in Fig. 4 in passing from **ZnP** to **1**. The Q band results from the S₀ → S₂₀, S₂₂ transitions computed around 540 nm with intensities ($f = 0.011$ and 0.006, respectively) similar to **ZnP**.

The formation of the positively charged species was furthermore investigated by spectrochemistry measurements through stepwise addition of FeCl₃. The UV-visible absorption spectra of the oxidized species of ZnP-2EDOTV-C₆₀ (**1**) in dichloromethane are plotted in Fig. 5, and those of **ZnP** and Ph-2EDOTV-C₆₀ (**6**) are displayed in Fig. S39 and S40 (ESI[‡]), respectively. For reference compounds **ZnP** and **6**, clear oxidation processes are observed, with the formation of exclusively one cation species, as corroborated by the observation of well-defined isosbestic points. In contrast, two oxidation processes can be seen for **1** (Fig. 5). After the initial addition of FeCl₃, the decrease of the band of the oligomer in the 450–500 nm range and the raise of a new band at 660 nm are observed. The new band is attributed to the formation of the cation ZnP-2EDOTV^{•+}-C₆₀ as compared with the spectrochemical oxidation of Ph-2EDOTV-C₆₀ (**6**) (Fig. S40, ESI[‡]). Theoretical calculations on the cation species of Ph-2EDOTV (**5**) and **6** further support this assignment by predicting an intense electronic excitation calculated at 621 nm ($f = 1.444$) for 5^{•+} and 653 nm ($f = 0.892$) for 6^{•+} (Fig. S37, ESI[‡]). Upon successive additions of FeCl₃, the band at 421 nm decreases in intensity and two new bands appear at 442 and 646 nm (Fig. 5). Similar bands are observed in the spectrochemical oxidation of **ZnP** at 438 and 635 nm (Fig. S39, ESI[‡]), which are theoretically predicted at 470 and 668 nm for ZnP^{•+} (Fig. S37, ESI[‡]), thus allowing the assignment of a radical located on the porphyrin moiety in the dication ZnP^{•+}-2EDOTV^{•+}-C₆₀, as also suggested by theoretical calculations (see above). These findings support the easiness to oxidize both the oligomer bridge and the porphyrin moiety, and the preference to stabilize the first radical cation on the conjugated bridge.

The photophysical properties of ZnP-2EDOTV-C₆₀ were studied by steady-state emission spectroscopy in toluene and benzonitrile (Fig. 6). The spectra of compounds Ph-2EDOTV (**5**) and Ph-2EDOTV-C₆₀ (**6**) are depicted, for comparison

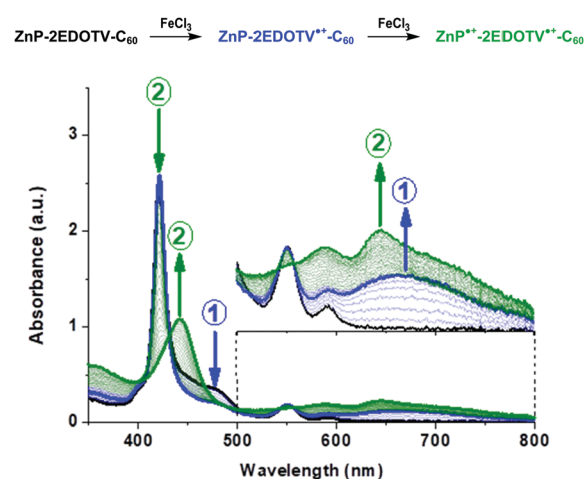


Fig. 5 UV-Vis absorption spectra of neutral (black), cation (blue) and dication (green) species of ZnP-2EDOTV-C₆₀ (**1**), in dichloromethane solution, generated upon successive additions of FeCl₃.

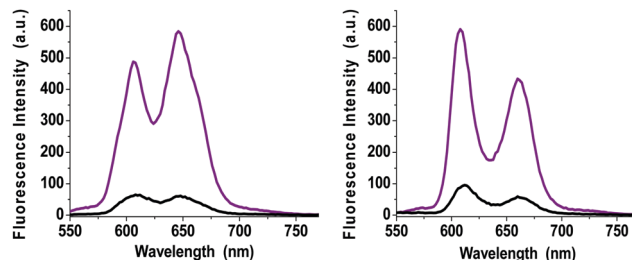


Fig. 6 Steady-state fluorescence spectra of **ZnP** (purple line) and **ZnP-2EDOTV-C₆₀** (**1**, black line) in toluene (left) and benzonitrile (right). $\lambda_{\text{exc}} = 426$ nm. Concentration 10^{-6} M.

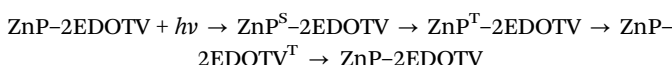
purposes, in Fig. S41 and S42 (ESI[‡]). For **1**, the excitation wavelength was 426 nm, where mainly the porphyrin absorbs. In both solvents, a quenching of the fluorescence of the porphyrin by around 90% is observed.

Transient absorption spectra

The photo-induced reactions of the compounds were studied using transient absorption (TA) spectroscopy, namely a femtosecond pump–probe method with excitation (pump) by a short 100 fs pulse and monitoring the change (probe) in the visible and near-infrared (NIR) regions. Most of the measurements were carried out using two excitation wavelengths, 495 and 560 nm, which allows to excite predominantly either 2EDOTV or **ZnP** chromophores, respectively.

The TA measurements on the Ph-2EDOTV (**5**) reference compound reveal that the singlet-excited state has a characteristic absorption band at 730 nm with a lifetime of 640 ps in PhCN (Fig. S43, ESI[‡]). The intersystem crossing efficiency of **5** is very low and the singlet state relaxes directly to the ground state leaving virtually no signal at delays longer than a few nanoseconds.

Excitation of dyad **4** (ZnP-2EDOTV) in toluene at 560 nm results in instant formation of the **ZnP** singlet-excited state, which decays to a state with a strong absorption band close to 610 nm, as presented in Fig. 7. This new band is formed with a time constant of 2.0 ± 0.2 ns, which is somewhat shorter than the lifetime of the **ZnP** singlet-excited state (2.4 ns). The spectrum of the state cannot be assigned neither to the **ZnP** triplet nor to the 2EDOTV singlet, and it is tentatively attributed to the 2EDOTV triplet, which can be reached in the case of rapid triplet–triplet energy transfer from **ZnP** to 2EDOTV, as will be discussed later. A tentative reaction scheme for ZnP-2EDOTV in toluene is



with the intersystem crossing ($\text{ZnP}^{\text{S}}\text{-2EDOTV} \rightarrow \text{ZnP}^{\text{T}}\text{-2EDOTV}$) and the energy transfer ($\text{ZnP}^{\text{T}}\text{-2EDOTV} \rightarrow \text{ZnP-2EDOTV}^{\text{T}}$) having roughly the same rates, so that population of $\text{ZnP}^{\text{T}}\text{-2EDOTV}$ is low at any delay time and this intermediate state is difficult to observe experimentally.

To ascertain the reaction scheme proposed above, the triplet excited states of ZnP-2EDOTV (**4**), **ZnP** and Ph-2EDOTV (**5**) were

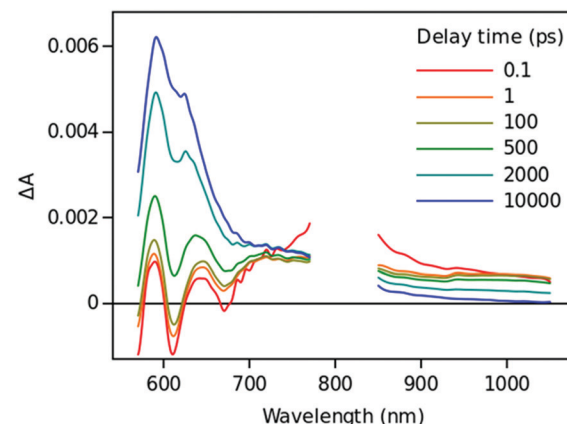


Fig. 7 Time-resolved TA spectra of ZnP-2EDOTV (**4**) in toluene excited at 560 nm.

calculated at the TD-B3LYP/(cc-pVDZ+LANL2DZ(Zn))+PCM level using toluene as solvent. The lowest-lying triplet state T_1 of ZnP-2EDOTV (**4**) is computed at 1.18 eV over the singlet ground state, and results from the HOMO \rightarrow LUMO excitation associated to the 2EDOTV moiety (see Fig. S27, ESI[‡]). This is in line with the T_1 state calculated for Ph-2EDOTV (**5**), which is predicted at a slightly higher energy (1.29 eV) due to a smaller π -conjugation compared to **4**. The unpaired-electron spin-density computed for **4** after fully optimizing the geometry of the T_1 state at the UB3LYP/(cc-pVDZ+LANL2DZ(Zn))+PCM(toluene) level spreads over the 2EDOTV bridge and is identical to that obtained for the T_1 of **5** (Fig. S44, ESI[‡]). The T_2 state of **4**, corresponding to the excitation of the **ZnP** moiety (HOMO-1 \rightarrow LUMO+1 excitation, Fig. S27, ESI[‡]), is calculated higher in energy at 1.57 eV, in good correlation with the lowest triplet state of **ZnP** (1.58 eV). Theoretical calculations therefore assign the lowest triplet of **4** to the $\text{ZnP-2EDOTV}^{\text{T}}$ species, and clearly support the reaction scheme described above with a downhill triplet–triplet energy transfer from **ZnP** to 2EDOTV.

In polar PhCN the TA response of the ZnP-2EDOTV dyad is more complex as presented in Fig. 8. Excitation at 560 nm results in an almost instant formation of a state with characteristic bleaching of the **ZnP** Q-bands (570 and 610 nm) and a deep at 670 nm which can be attributed to the **ZnP** stimulated emission typical of porphyrins. At the same time, a strong absorption band is formed in the red side with maximum close to 770 nm, which resembles the singlet state of Ph-2EDOTV. However, this band is much broader, exhibits a red shoulder extended to the NIR and differs significantly from the band of the Ph-2EDOTV singlet state (Fig. S43, ESI[‡]).

One can notice that HOMO energies of **ZnP** and Ph-2EDOTV are very close to each other (Fig. 2 and Fig. S28 in the ESI[‡]), and that the HOMO of Ph-2EDOTV is only marginally higher than that of **ZnP**. In the singlet-excited state ZnP^{S} , the HOMO of **ZnP** has one vacancy, which makes possible the hole transfer from the **ZnP** HOMO to the 2EDOTV HOMO, thus generating the charge-separated state $\text{ZnP}^{\text{-}}\text{-2EDOTV}^{\text{+}}$. However, we do not observe pure **ZnP** singlet-excited state with the time resolution of our instrument, roughly 200 fs. At 0.2 ps delay, the state with

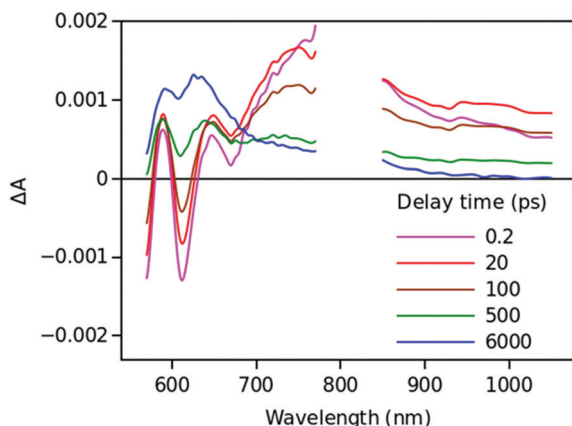
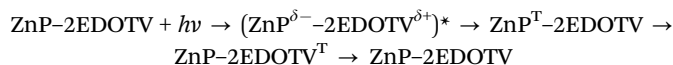


Fig. 8 Time-resolved TA spectra of ZnP-2EDOTV (**4**) in PhCN excited at 560 nm.

features of both **ZnP** and 2EDOTV excited states is seen instead. This is typical of the exciplex-like state, which involves both chromophores and is characterized by a partial electron density transfer, $(\text{ZnP}^{\delta-} \text{--} 2\text{EDOTV}^{\delta+})^*$. Furthermore, one can speculate that this state is in equilibrium with the **ZnP**^S locally excited state as indicated by a deep at 670 nm, which is typical of a **ZnP** stimulated emission feature. The global data fit shows that this state decays with time constant 160 ± 20 ps (Fig. S45, ESI‡). The relaxation is mainly seen as: (1) disappearance of the broad band absorption in the range from 700 nm to the NIR and (2) disappearance of the deep at 670 nm. The time-resolved spectrum at 500 ps delay is representative of this new intermediate state, it holds only features of the **ZnP** chromophore and is tentatively attributed to the **ZnP** triplet state. Unlike the triplet state of the isolated **ZnP** chromophore, the lifetime of this triplet in ZnP-2EDOTV is relatively short, and it does not relax directly to the ground state but decays with a time constant of 1.9 ± 0.6 ns to a long-lived state with the spectrum represented by the time-resolved spectrum at 6000 ps in Fig. 8. This spectrum can be assigned to the triplet state of 2EDOTV and the whole reaction scheme is



Switching the pump wavelength to 495 nm, which excites predominantly the 2EDOTV chromophore in the ZnP-2EDOTV dyad, does not change the TA response observed with 200 fs time resolution as presented in Fig. S43 and S45 (ESI‡). One can expect efficient energy transfer from the excited singlet state of 2EDOTV to **ZnP**, but the fact that we do not time-resolve this process can be interpreted in favor of a Dexter energy-transfer mechanism and a relatively strong electronic coupling between the two chromophores. This, the strong electronic coupling, explains relatively fast triplet state energy transfer taking approximately 2 ns.

The main difference between the photophysics in non-polar toluene and polar PhCN is the formation of the $(\text{ZnP}^{\delta-} \text{--} 2\text{EDOTV}^{\delta+})^*$ state, which is not observed in toluene. Since this state is

characterized by a charge shift, its energy depends on the solvent polarity and it can be speculated that in toluene it is higher than that of the localized singlet state $\text{ZnP}^S\text{-2EDOTV}$, and thus it is not observed. However, the energy transfer is rather insensitive to the solvent polarity and should be roughly the same in toluene and PhCN, or close to 2 ns, which is faster than intersystem crossing of **ZnP**. As the result, the triplet state of the **ZnP** chromophore is almost invisible in toluene as its relaxation rate (*via* energy transfer) is faster than the formation rate.

Excitation of the Ph-2EDOTV-C₆₀ (**6**) dyad at 420 nm results in an instant formation of the 2EDOTV^S excited state, which relaxes quickly by yielding the 2EDOTV⁺-C₆₀⁻ CS state followed by a rapid charge recombination. The three lowest-lying singlet states calculated for **6** below 1.55 eV actually correspond to 2EDOTV⁺-C₆₀⁻ CS states resulting from an electron transfer from the HOMO, located on the Ph-2EDOTV moiety, to the LUMO+1, LUMO+2 and LUMO+3 spreading over the C₆₀ cage (Fig. S27, ESI‡). Fig. 9 presents the TA time-resolved spectra of the dyad in toluene. The spectrum at 0.2 ps delay time has one clear absorption maximum at 710 nm, which matches well that of the singlet-excited state of the reference compound Ph-2EDOTV (Fig. S43, ESI‡). Within few ps, a new absorption band develops at 1080 nm, which can be attributed to the fullerene anion C₆₀⁻. At the same time, the band at 710 nm shifts to the blue (maximum close to 690 nm) and increases in intensity gradually. This band can be attributed to the cation 2EDOTV⁺, and correlates with the intense transition theoretically predicted for **6**⁺ at 653 nm (*f* = 0.892). The CS state recombines with a time constant of 160 ± 10 ps according to the global fit results.

In PhCN, both charge separation and recombination are faster, which is typical for polar solvents as compared to non-polar (Fig. S48, ESI‡). The estimated time constants for the charge separation and recombination in PhCN are 2.4 ± 0.3 and 4.1 ± 0.4 ps, respectively. This behavior agrees with the energy diagram presented in Fig. 2, showing a rather large driving force for the CS and a marginally higher driving force for the recombination, which makes the recombination a fast process as well.

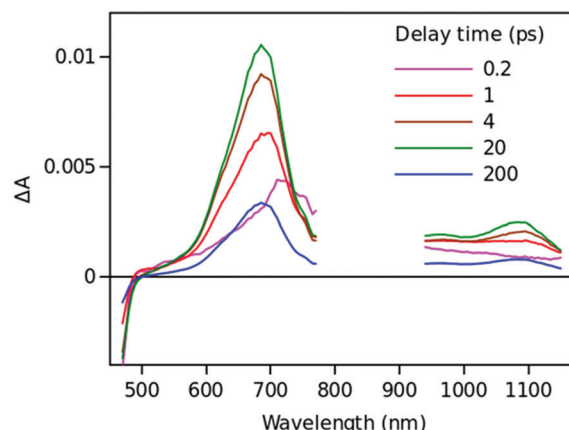


Fig. 9 Time resolved TA spectra of Ph-2EDOTV-C₆₀ (**6**) in toluene excited at 420 nm.

Time resolved TA spectra of the triad ZnP-2EDOTV-C₆₀ (**1**) in toluene and PhCN obtained with excitation at 550 nm are presented in Fig. 10 and 11, respectively. The main difference between the responses in these two solvents is the time scale, the spectra shapes having many common features.

At a short delay time (0.2 ps in Fig. 10), the time-resolved spectra resemble the singlet state ZnP^S most closely. Further evolution of the spectrum is associated with the buildup of two broad absorption bands at roughly 720 and 970 nm and the disappearance of the deep at 670 nm. Characteristic spectra of this intermediate can be seen at roughly 6 and 40 ps delay time in PhCN and toluene, respectively. At least in PhCN, this spectrum resembles that of the ZnP⁺-C₆₀⁻ charge-separated state observed for numerous porphyrin-fullerene dyads though typical position of the fullerene anion radical band is at 1040 nm but not at 970 nm. However, in porphyrin-fullerene dyads with close proximity of the donor and acceptor moieties, an exciplex-like state is formed with the electron density spreading over both chromophores, ZnP^{δ+}-C₆₀^{δ-}, instead of being localized completely on the acceptor. In this case, the fullerene anion-type band is shifted to 950–1000 nm and broadened.^{29,30} The exciplex-like state can be expected in the case of ZnP-2EDOTV-C₆₀ compound as well, and it explains the spectrum shape in the NIR part.

There is a noticeable difference in the visible spectra between the sample in toluene and PhCN. The characteristic feature of the ZnP chromophore is the bleaching of the Q-band at 590 nm. This feature almost vanishes at 40 ps delay in toluene. At the same time, the rest of the spectrum resembles the spectrum of Ph-2EDOTV-C₆₀ at similar delay time (20 ps in Fig. 9), which was assigned to the Ph-2EDOTV⁺-C₆₀⁻ CS state, though there is a perceptible difference between the spectra in the NIR. Therefore, the formation of an exciplex-like state between Ph-2EDOTV and C₆₀, Ph-2EDOTV^{δ+}-C₆₀^{δ-}, can be assumed. As mentioned above, the energies of the HOMO levels of ZnP and 2EDOTV are very close to each other and the energy difference between the states ZnP⁺-C₆₀⁻ and 2EDOTV⁺-C₆₀⁻, or ZnP^{δ+}-C₆₀^{δ-} and 2EDOTV^{δ+}-C₆₀^{δ-}, must

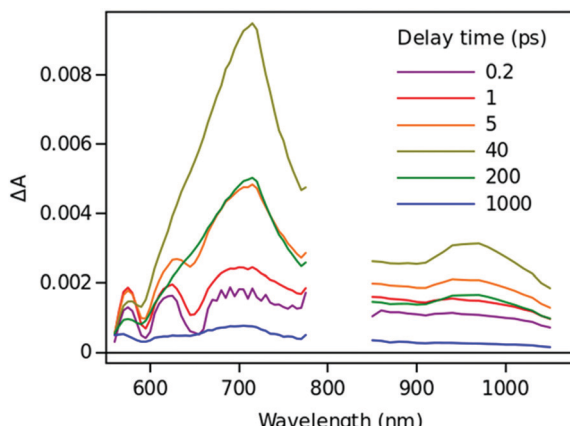


Fig. 10 Time resolved TA spectra of ZnP-2EDOTV-C₆₀ (**1**) in toluene excited at 550 nm.

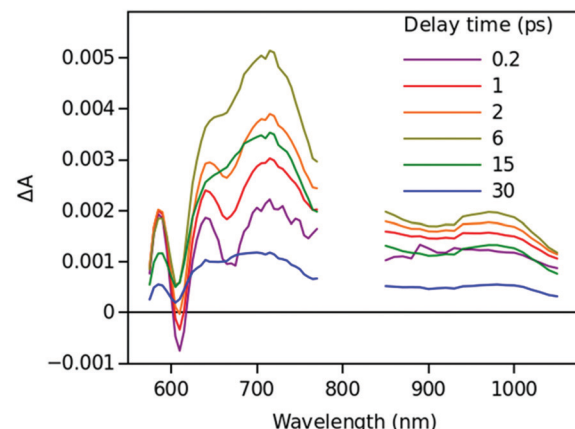
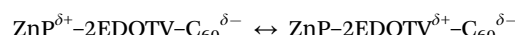
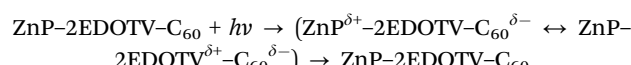


Fig. 11 Time resolved TA spectra of ZnP-2EDOTV-C₆₀ (**1**) in PhCN excited at 550 nm.

also be relatively low. This opens the possibility for the equilibrium



which is achieved through a hole transfer between the HOMOs of ZnP and 2EDOTV chromophores. Then, the second difference between ZnP-2EDOTV-C₆₀ in toluene and PhCN is the shift in the equilibrium between these two exciplex states. In PhCN the equilibrium is more on the side of ZnP^{δ+}-C₆₀^{δ-}, and in toluene is on the side of 2EDOTV^{δ+}-C₆₀^{δ-}. The reaction scheme is



Only two time constants are available from the TA measurements, 12 and 210 ps in toluene and 5 and 11 ps in PhCN. The first one corresponds to the charge separation and the establishment of the equilibrium, and the second one governs the relaxation to the ground state. The equilibrium rate constant is high in both cases and cannot be extracted from the TA measurements.

The charge transfer distance for the state ZnP^{δ+}-2EDOTV-C₆₀^{δ-} can be estimated to be ~2.45 nm and it is achieved in both solvents in a few picoseconds time. This is a long distance in the molecular scale, but this CS happens without the energy loss typically found for the more traditional design of molecular systems consisting of multiple donors and acceptors and implementing a multi-step ET,³¹ in which case the energy is lost at each sequential ET step. The long CS distance in ZnP-2EDOTV-C₆₀ compound is achieved through enhanced electronic communication between the donor (ZnP) and the acceptor (C₆₀) via the conjugated bridge (2EDOTV). This approach has been used in a number of D-B-A system designs leading to a one-step, long-distance CS via a molecular wire connecting the donor and acceptor.^{32,33} The time constants of the CS here reported are comparable to those systems with similar DA separation distance. A disadvantage of the DA electronic coupling enhanced by the molecular wire strategy

is the accelerated charge recombination as well, which is also observed in the case of the ZnP-2EDOTV-C₆₀ compound.

Conclusions

The design goal of the ZnP-2EDOTV-C₆₀ triad is to provide relatively high electronic coupling between the electron donor (ZnP) and electron acceptor (C₆₀) moieties through the π -conjugated bridge (2EDOTV), which was expected to accelerate the photoinduced electron transfer. The porphyrin-C₆₀ centroid-centroid distance is estimated to be 24.5 Å for this compound and even in non-polar toluene the CS is as fast as 12 ps, and the goal of the fast CS is achieved. The observed equilibrium of hole shifting between ZnP and 2EDOTV in the triad (ZnP^{δ+}-2EDOTV-C₆₀^{δ-} ↔ ZnP-2EDOTV^{δ+}-C₆₀^{δ-}) can be considered as a drawback. However, we can envision that this problem can be solved by a more careful selection of the component energies. In particular, a conjugated bridge with a HOMO level lower in energy than that of the donor moiety in the D-B-A triad should help to avoid undesired intermediate states and to achieve a more complete charge separation with slower charge recombination.

Conflicts of interest

The authors declare no competing financial interests.

Acknowledgements

The authors thank the Spanish Ministry of Science and Innovation (MICINN) (PID2019-105049RB-I00, PGC2018-099568-B-I00, RED2018-102815-T and Unidad de Excelencia María de Maeztu CEX2019-000919-M), the Junta de Comunidades de Castilla-La Mancha and European Social Fund (SBPLY/17/180501/000254), the Generalitat Valenciana (PROMETEO/2020/077 and SEJ/2018/035) and European FEDER funds (PGC2018-099568-B-I00) for financial support. J. A. acknowledges the MICINN for his "Ramón-y-Cajal" Fellowship (RyC-2017-23500).

References

- 1 R. E. Blankenship, *Molecular Mechanisms of Photosynthesis*, Wiley-Blackwell, Chichester, 2001.
- 2 I. McConnell, G. Li and G. W. Brudvig, *Chem. Biol.*, 2010, **17**, 434–447.
- 3 M. R. Wasielewski, *Chem. Rev.*, 1992, **92**, 435–461.
- 4 D. Gust, T. A. Moore and A. L. Moore, *Acc. Chem. Res.*, 2001, **34**, 40–48.
- 5 M. E. El-Khoudy, E. El-Moshnawy and S. Fukuzumi, *J. Photochem. Photobiol. C*, 2017, **31**, 36–83.
- 6 M. Mamedov, Govindjee, V. Nadtochenko and A. Semenov, *Photosynth. Res.*, 2015, **125**, 51–63.
- 7 A. A. Strelnikov, A. S. Konev, O. V. Levin, A. F. Khlebnikov, A. Iwasaki, K. Yamanouchi and N. V. Tkachenko, *J. Phys. Chem. B*, 2020, **124**, 10899–10912.
- 8 C. Schubert, J. T. Margraf, T. Clark and D. M. Guldi, *Chem. Soc. Rev.*, 2002, **44**(2015), 988–998.
- 9 T. Higashino, T. Yamada, M. Yamamoto, A. Furube, N. V. Tkachenko, T. Miura, Y. Kobori, R. Jono, K. Yamashita and H. Imahori, *Angew. Chem., Int. Ed.*, 2016, **55**, 629–633.
- 10 F. Oswald, D.-M. Shafiqul-Islam, Y. Araki, V. Troiani, R. Caballero, P. de la Cruz, O. Ito and F. Langa, *Chem. Commun.*, 2007, 4498–4500.
- 11 H. B. Gray and J. R. Winkler, *Proc. Natl. Acad. Sci. U. S. A.*, 2005, **102**, 3534–3539.
- 12 H. Oevering, M. N. Paddon-Row, M. Heppener, A. M. Oliver, E. Cotsaris, J. W. Verhoeven and N. S. Hush, *J. Am. Chem. Soc.*, 1987, **109**, 3258–3269.
- 13 O. S. Wenger, *Acc. Chem. Res.*, 2011, **44**, 25–35.
- 14 J. Sukegawa, C. Schubert, X. Zhu, H. Tsuji, D. M. Guldi and E. Nakamura, *Nat. Chem.*, 2014, **6**, 899–905.
- 15 D. M. Guldi, *Chem. Soc. Rev.*, 2002, **31**, 22–36.
- 16 H. Imahori and Y. Sakata, *Adv. Mater.*, 1997, **9**, 537–546.
- 17 J. M. Tour, *Acc. Chem. Res.*, 2000, **33**, 791–804.
- 18 G. de la Torre, F. Giacalone, J. L. Segura, N. Martín and D. M. Guldi, *Chem. – Eur. J.*, 2005, **11**, 1267–1280.
- 19 A. Lembo, P. Tagliatesta, D. M. Guldi, M. Wielopolski and M. Nuccetelli, *J. Phys. Chem. A*, 2009, **113**(9), 1779–1793.
- 20 M. Wielopolski, G. de Miguel Rojas, C. van del Pol, L. Brinkhaus, G. Katsukis, M. R. Bryce, T. Clark and D. M. Guldi, *ACS Nano*, 2010, **4**(11), 6449–6462.
- 21 J. Ikemoto, K. Takimiya, Y. Aso, T. Otsubo, M. Fujitsuka and O. Ito, *Org. Lett.*, 2002, **4**, 309–311.
- 22 F. Oswald, D.-M. Shafiqul Islam, M. E. El-Khoudy, Y. Araki, R. Caballero, P. de la Cruz, O. Ito and F. Langa, *Phys. Chem. Chem. Phys.*, 2014, **16**, 2443–2451.
- 23 C. Franco, P. Mayorga Berriezo, V. Lloveras, R. Caballero, I. Alcón, S. T. Bromley, M. Mas-Torrent, F. Langa, J. T. López-Navarrete, C. Rovira, J. Casado and J. Veciana, *J. Am. Chem. Soc.*, 2017, **139**, 686–692.
- 24 P. Mayorga-Burrezo, B. Pelado, R. Ponce-Ortiz, P. De la Cruz, J. T. López-Navarrete, F. Langa and J. Casado, *Chem. – Eur. J.*, 2015, **21**, 1713–1725.
- 25 A. Balakumar, A. B. Lysenko, C. Carcel, V. L. Malinovskii, D. T. Gryko, K.-H. Schweikart, R. S. Loewe, A. A. Yasseri, Z. Liu, D. F. Bocian and J. S. Lindsey, *J. Org. Chem.*, 2004, **69**, 1453–1460.
- 26 B. Pelado, P. De La Cruz, V. González-Pedro, E. M. Barea and F. Langa, *Tetrahedron Lett.*, 2012, **53**, 6665–6669.
- 27 C. Martineau, P. Blanchard, D. Rondeau, J. Delaunay and J. Roncali, *Adv. Mater.*, 2002, **14**, 283–287.
- 28 The energy of each radical ion pair (RIP) is estimated from electrochemical measurements by applying the following equation ($E_{\text{RIP}} = E_{\text{ox}} - E_{\text{red}}$). E_{ox} is 0.21 V for the radical ion pair ZnP-2EDOTV⁺-C₆₀^{•-} and 0.28 V for ZnP^{•+}-2EDOTV-C₆₀^{•-}.
- 29 N. V. Tkachenko, H. Lemmetyinen, J. Sonoda, K. Ohkubo, T. Sato, H. Imahori and S. Fukuzumi, *J. Phys. Chem. A*, 2003, **107**, 8834–8839.

30 V. Chukharev, N. V. Tkachenko, A. Efimov, D. M. Guldi, A. Hirsch, M. Scheloske and H. Lemmetyinen, *J. Phys. Chem. B*, 2004, **108**, 16377–16385.

31 H. Imahori, Y. Sekiguchi, Y. Kashiwagi, T. Sato, Y. Araki, O. Ito, H. Yamada and S. Fukuzumi, *Chem. – Eur. J.*, 2004, **10**, 3184–3196.

32 W. B. Davis, M. A. Ratner and M. R. Wasielewski, *J. Am. Chem. Soc.*, 2001, **123**, 7877–7886.

33 J. Sukegawa, C. Schubert, X. Zhu, H. Tsuji, D. M. Guldi and E. Nakamura, *Nat. Chem.*, 2014, **6**, 899–905.